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NEWS	1		Web Page URLs for STN Seminar Schedule - N. America
NEWS	2		"Ask CAS" for self-help around the clock
NEWS	3	FEB 25	CA/CAPLUS - Russian Agency for Patents and Trademarks (ROSPATENT) added to list of core patent offices covered
NEWS	4	FEB 28	PATDPAFULL - New display fields provide for legal status data from INPADOC
NEWS	5	FEB 28	BABS - Current-awareness alerts (SDIs) available
NEWS	6	FEB 28	MEDLINE/LMEDLINE reloaded
NEWS	7	MAR 02	GBFULL: New full-text patent database on STN
NEWS	8	MAR 03	REGISTRY/ZREGISTRY - Sequence annotations enhanced
NEWS	9	MAR 03	MEDLINE file segment of TOXCENTER reloaded
NEWS	10	MAR 22	KOREAPAT now updated monthly; patent information enhanced
NEWS	11	MAR 22	Original IDE display format returns to REGISTRY/ZREGISTRY
NEWS	12	MAR 22	PATDPASPC - New patent database available
NEWS	13	MAR 22	REGISTRY/ZREGISTRY enhanced with experimental property tags
NEWS	14	APR 04	EPFULL enhanced with additional patent information and new fields
NEWS	15	APR 04	EMBASE - Database reloaded and enhanced
NEWS	16	APR 18	New CAS Information Use Policies available online
NEWS	17	APR 25	Patent searching, including current-awareness alerts (SDIs), based on application date in CA/CAPLUS and USPATFULL/USPAT2 may be affected by a change in filing date for U.S. applications.
NEWS	18	APR 28	Improved searching of U.S. Patent Classifications for U.S. patent records in CA/CAPLUS
NEWS EXPRESS			JANUARY 10 CURRENT WINDOWS VERSION IS V7.01a, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), AND CURRENT DISCOVER FILE IS DATED 10 JANUARY 2005
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FILE 'HOME' ENTERED AT 14:23:43 ON 09 MAY 2005

=> index bioscience

FILE 'DRUGMONOG' ACCESS NOT AUTHORIZED

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ENTRY	SESSION
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75 FILES IN THE FILE LIST IN STNINDEX

Enter SET DETAIL ON to see search term postings or to view search error messages that display as 0* with SET DETAIL OFF.

=> s preconditioning

370	FILE ADISCTI
18	FILE ADISINSIGHT
14	FILE ADISNEWS
275	FILE AGRICOLA
70	FILE ANABSTR
14	FILE ANTE
27	FILE AQUALINE
140	FILE AQUASCI
121	FILE BIOBUSINESS
4	FILE BIOCOMMERCE
131	FILE BIOENG
6800	FILE BIOSIS
109	FILE BIOTECHABS
109	FILE BIOTECHDS
412	FILE BIOTECHNO
657	FILE CABA
315	FILE CANCERLIT
5254	FILE CAPLUS
54	FILE CEABA-VTB
15	FILE CIN
267	FILE CONFSCI
15	FILE CROPB
59	FILE CROPU
5	FILE DDFB
869	FILE DDFU
76	FILE DGENE
553	FILE DISSABS
5	FILE DRUGB
994	FILE DRUGU
82	FILE EMBAL
4241	FILE EMBASE
2600	FILE ESBIODASE
206	FILE FEDRIP
65	FILE FROSTI
98	FILE FSTA
61	FILE GENBANK
9	FILE HEALSAFE
1201	FILE IFIPAT
3	FILE IMSDRUGNEWS
4	FILE IMSRESEARCH
711	FILE JICST-EPLUS
7	FILE KOSMET
526	FILE LIFESCI
6	FILE MEDICONF

4881 FILE MEDLINE
 24 FILE NIOSHTIC
 583 FILE NTIS
 52 FILES SEARCHED...
 75 FILE OCEAN
 5144 FILE PASCAL
 2 FILE PHAR
 3 FILE PHARMAML
 18 FILE PHIN
 306 FILE PROMT
 6 FILE PROUSDDR
 9 FILE RDISCLOSURE
 8243 FILE SCISEARCH
 2045 FILE TOXCENTER
 4450 FILE USPATFULL
 361 FILE USPAT2
 8 FILE VETB
 14 FILE VETU
 99 FILE WATER
 718 FILE WPIDS
 3 FILE WPIFV
 718 FILE WPINDEX

65 FILES HAVE ONE OR MORE ANSWERS, 75 FILES SEARCHED IN STNINDEX

L1 QUE PRECONDITIONING

=> d rank

F1	8243	SCISEARCH
F2	6800	BIOSIS
F3	5254	CAPLUS
F4	5144	PASCAL
F5	4881	MEDLINE
F6	4450	USPATFULL
F7	4241	EMBASE
F8	2600	ESBIOBASE
F9	2045	TOXCENTER
F10	1201	IFIPAT
F11	994	DRUGU
F12	869	DDFU
F13	718	WPIDS
F14	718	WPINDEX
F15	711	JICST-EPLUS
F16	657	CABA
F17	583	NTIS
F18	553	DISSABS
F19	526	LIFESCI
F20	412	BIOTECHNO
F21	370	ADISCTI
F22	361	USPAT2
F23	315	CANCERLIT
F24	306	PROMT
F25	275	AGRICOLA
F26	267	CONFSCI
F27	206	FEDRIP
F28	140	AQUASCI
F29	131	BIOENG
F30	121	BIOBUSINESS
F31	109	BIOTECHABS
F32	109	BIOTECHDS
F33	99	WATER
F34	98	FSTA
F35	82	EMBAL

F36	76	DGENE
F37	75	OCEAN
F38	70	ANABSTR
F39	65	FROSTI
F40	61	GENBANK
F41	59	CROPU
F42	54	CEABA-VTB
F43	27	AQUALINE
F44	24	NIOSHTIC
F45	18	ADISINSIGHT
F46	18	PHIN
F47	15	CIN
F48	15	CROPB
F49	14	ADISNEWS
F50	14	ANTE
F51	14	VETU
F52	9	HEALSAFE
F53	9	RDISCLOSURE
F54	8	VETB
F55	7	KOSMET
F56	6	MEDICONF
F57	6	PROUSDDR
F58	5	DDFB
F59	5	DRUGB
F60	4	BIOCOMMERCE
F61	4	IMSRESEARCH
F62	3	IMSDRUGNEWS
F63	3	PHARMAML
F64	3	WPIFV
F65	2	PHAR

=> file f1, f2, f3, f4, f5, f6, f7, f8, f9
COST IN U.S. DOLLARS

	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	1.18	1.39

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=> s preconditioning (p) mitochondria
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
FIELD CODE - 'AND' OPERATOR ASSUMED 'DITONING (P) MITOCHOND'
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
FIELD CODE - 'AND' OPERATOR ASSUMED 'DITONING (P) MITOCHOND'
L2 1261 PRECONDITIONING (P) MITOCHONDRIA

=> dup rem l2
PROCESSING COMPLETED FOR L2
L3 508 DUP REM L2 (753 DUPLICATES REMOVED)

=> s L3 AND (isocitrate(w)dehydrogenase(w)NAD(w)alpha(w)subunit OR
succinyl(w)CoA(w)ligase OR ATP(w)synthase OR dihyrolipoamide(w)succinyltransferase
OR ubiquinol(w)cytochrome(w)c(w)oxidoreductase OR
NADH(w)ubiquinone(w)oxidoreductase)
6 FILES SEARCHED...

L4 15 L3 AND (ISOCITRATE(W) DEHYDROGENASE(W) NAD(W) ALPHA(W) SUBUNIT
OR SUCCINYL(W) COA(W) LIGASE OR ATP(W) SYNTHASE OR DIHYROLIPOAMI
DE(W) SUCCINYLTRANSFERASE OR UBIQUINOL(W) CYTOCHROME(W) C(W)
OXIDOREDUCTASE OR NADH(W) UBIQUINONE(W) OXIDOREDUCTASE)

=> d l4 ibib ti abs 1-15

L4 ANSWER 1 OF 15 SCISEARCH COPYRIGHT (c) 2005 The Thomson Corporation on
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ACCESSION NUMBER: 2004:489036 SCISEARCH

THE GENUINE ARTICLE: 822TJ

TITLE: Diazoxide affects the IF1 inhibitor protein binding to F-1
sector of beef heart F(0)F(1)ATPsynthase

AUTHOR: Contessi S; Metelli G; Mavelli I; Lippe G (Reprint)

CORPORATE SOURCE: Univ Udine, Dept Biomed Sci & Technol, Ple Kolbe 4,
I-33100 Udine, Italy (Reprint); Univ Udine, Dept Biomed
Sci & Technol, I-33100 Udine, Italy; Univ Udine, MATI Ctr
Excellence, I-33100 Udine, Italy

COUNTRY OF AUTHOR: Italy

SOURCE: BIOCHEMICAL PHARMACOLOGY, (15 MAY 2004) Vol. 67, No. 10,
pp. 1843-1851.

Publisher: PERGAMON-ELSEVIER SCIENCE LTD, THE BOULEVARD,
LANGFORD LANE, KIDLINGTON, OXFORD OX5 1GB, ENGLAND.

ISSN: 0006-2952.

DOCUMENT TYPE: Article; Journal

LANGUAGE: English

REFERENCE COUNT: 39

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

TI Diazoxide affects the IF1 inhibitor protein binding to F-1 sector of beef
heart F(0)F(1)ATPsynthase

AB Diazoxide, a selective opener of the mitochondrial ATP-sensitive K⁺
channel (mitoK(ATP)), has been reported to enhance F(0)F(1)ATP-
synthase inhibition during ischemia, but the underlying mechanisms
are still unclear. Here, we demonstrate that diazoxide directly interacts
with the F-1 sector of beef heart F(0)F(1)ATPsynthase markedly promoting
the binding of the inhibitor protein (IF1) to beta subunit. More
specifically, the treatment of soluble F-1 with one equivalent of
diazoxide was sufficient to decrease the K_d of IF1-F-1 complex at low pH.
Such effect was revealed only on the cycling enzyme, while no effect was
observed in the absence of Mg-ATP. However, diazoxide binding occurred
independently from the catalysis, as shown by the structural changes
induced by the drug in not catalytically active F-1 and revealed by CD

spectra. In addition, kinetic analysis of ATP hydrolysis demonstrated that diazoxide exerts a stabilising role on Mg-ADP bound in the catalytic site of the beta subunit adopting the tight conformation (beta(DP)). In accordance, a stabilising effect of Mg-ADP at the nucleotide binding domain (NBD) has been reported also for K-ATP channel. These results suggest that diazoxide binds to beta subunit at NBD, which is highly conserved in the ATP-binding cassette protein family, thus inducing nucleotide stabilisation and favouring F-1 conformation suitable for IF₁ binding. Finally, diazoxide also increased IF₁ binding to membrane bound F₁, while it did not influence the energisation-dependent IF₁ release. As IF₁ binding mediates the F(0)F(1)ATP synthase inhibition, we suggest that such mechanism may contribute to cardioprotection during ischemia. (C) 2004 Elsevier Inc. All rights reserved.

L4 ANSWER 2 OF 15 SCISEARCH COPYRIGHT (c) 2005 The Thomson Corporation on STN
 ACCESSION NUMBER: 2003:972343 SCISEARCH
 THE GENUINE ARTICLE: 739VH
 TITLE: ATP-sensitive K⁺ channels in renal mitochondria
 AUTHOR: Cancherini D V; Trabuco L G; Reboucas N A; Kowaltowski A J (Reprint)
 CORPORATE SOURCE: Av Prof Lineu Prestes 748, Cidade Univ, BR-05508900 Sao Paulo, Brazil (Reprint); Univ Sao Paulo, Dept Fisiol & Biofis, Inst Ciencias Biomed, BR-05508900 Sao Paulo, Brazil; Univ Sao Paulo, Dept Bioquim, Inst Quim, BR-05508900 Sao Paulo, Brazil
 COUNTRY OF AUTHOR: Brazil
 SOURCE: AMERICAN JOURNAL OF PHYSIOLOGY-RENAL PHYSIOLOGY, (DEC 2003 Vol. 285, No. 6, pp. F1291-F1296. Publisher: AMER PHYSIOLOGICAL SOC, 9650 ROCKVILLE PIKE, BETHESDA, MD 20814 USA. ISSN: 0363-6127.
)
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
 REFERENCE COUNT: 38

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

TI ATP-sensitive K⁺ channels in renal mitochondria
 AB Isolated kidney mitochondria swell when incubated in hyposmotic solutions containing K⁺ salts in a manner inhibited by ATP, ADP, 5-hydroxydecanoate, and glibenclamide and stimulated by GTP and diazoxide. These results suggest the existence of ATP-sensitive K⁺ channels in these mitochondria, similar to those previously described in heart, liver, and brain. Renal mitochondrial ATP-sensitive K⁺ uptake rates are similar to 140 nmol . min⁽⁻¹⁾ . mg protein⁽⁻¹⁾. This K⁺ transport results in a slight increase in respiration and decrease in the inner membrane potential. In addition, the activation of ATP-inhibited K⁺ uptake using diazoxide leads to a decrease of ATP hydrolysis through the reverse activity of the F₀F₁ **ATP synthase** when respiration is inhibited. In conclusion, we characterize an ATP-sensitive K⁺ transport pathway in kidney mitochondria that affects volume, respiration, and membrane potential and may have a role in the prevention of mitochondrial ATP hydrolysis.

L4 ANSWER 3 OF 15 SCISEARCH COPYRIGHT (c) 2005 The Thomson Corporation on STN
 ACCESSION NUMBER: 2000:931072 SCISEARCH
 THE GENUINE ARTICLE: 379KX
 TITLE: Myocardial ischemic preconditioning and mitochondrial F₁F₀-ATPase activity
 AUTHOR: Bosetti F (Reprint); Yu G Y; Zucchi R; Ronca Testoni S; Solaini G
 CORPORATE SOURCE: NIA, SECT BRAIN PHYSIOL & METAB, NIH BLDG 10, ROOM 6N202, 9000 ROCKVILLE PIKE, BETHESDA, MD 20892 (Reprint); SCUOLA

SUPER STUDI UNIV & PERFEZIONAMENTO S ANNA, PISA, ITALY;
UNIV PISA, SEZ CHIM & BIOCHIM MED, DIPARTIMENTO SCI UOMO &
AMBIENTE, I-56100 PISA, ITALY

COUNTRY OF AUTHOR: USA; ITALY

SOURCE: MOLECULAR AND CELLULAR BIOCHEMISTRY, (DEC 2000) Vol. 215,
No. 1-2, pp. 31-37.
Publisher: KLUWER ACADEMIC PUBL, SPUIBOULEVARD 50, PO BOX
17, 3300 AA DORDRECHT, NETHERLANDS.
ISSN: 0300-8177.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: LIFE

LANGUAGE: English

REFERENCE COUNT: 39

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

TI Myocardial ischemic preconditioning and mitochondrial FlF₀-ATPase activity

AB A short period of ischemia followed by reperfusion (ischemic **preconditioning**) is known to trigger mechanisms that contribute to the prevention of ATP depletion. In ischemic conditions, most of the ATP hydrolysis can be attributed to mitochondrial FlF₀-ATPase (**ATP synthase**). The purpose of the present study was to examine the effect of myocardial ischemic **preconditioning** on the kinetics of ATP hydrolysis by FlF₀-ATPase. **Preconditioning** was accomplished by three 3-min periods of global ischemia separated by 3 min of reperfusion. Steady state ATP hydrolysis rates in both control and preconditioned **mitochondria** were not significantly different. This suggests that a large influence of the enzyme on the **preconditioning** mechanism may be excluded. However, the time required by the reaction to reach the steady state rate was increased in the preconditioned group before sustained ischemia, and it was even more enhanced in the first 5 min of reperfusion (101 +/- 3.0 sec in preconditioned vs. 83.4 +/- 4.4 sec in controls, p < 0.05). These results suggest that this transient increase in activation time may contribute to the cardioprotection by slowing the ATP depletion in the very critical early phase of post-ischemic reperfusion.

L4 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:692747 CAPLUS

DOCUMENT NUMBER: 141:241156

TITLE: Ischemic preconditioning exaggerates cardiac damage in PKC- δ null mice

AUTHOR(S): Mayr, Manuel; Metzler, Bernhard; Chung, Yuen-Li; McGregor, Emma; Mayr, Ursula; Troy, Helen; Hu, Yanhua; Leitges, Michael; Pachinger, Otmar; Griffiths, John R.; Dunn, Michael J.; Xu, Qingbo

CORPORATE SOURCE: Department of Cardiac and Vascular Sciences, St. George's Hospital Medical School, London, SW17 0RE, UK

SOURCE: American Journal of Physiology (2004), 287(2, Pt. 2), H946-H956
CODEN: AJPHAP; ISSN: 0002-9513

PUBLISHER: American Physiological Society

DOCUMENT TYPE: Journal

LANGUAGE: English

TI Ischemic preconditioning exaggerates cardiac damage in PKC- δ null mice

AB Ischemic preconditioning confers cardiac protection during subsequent ischemia-reperfusion, in which protein kinase C (PKC) is believed to play an essential role, but controversial data exist concerning the PKC- δ isoform. In an accompanying study, the authors described metabolic changes in PKC- δ knockout mice. The authors now explore their effect on early preconditioning. Both PKC- δ ^{-/-} and PKC- δ ^{+/-} mice underwent three cycles of 5-min left descending artery occlusion/5-min reperfusion, followed by 30-min occlusion and 2-h reperfusion. Unexpectedly, preconditioning exaggerated

ischemia-reperfusion injury in PKC- δ ^{-/-} mice. Whereas ischemic preconditioning increased superoxide anion production in PKC- δ ^{+/+} hearts, no increase in reactive oxygen species was observed in PKC- δ ^{-/-} hearts. Proteomic anal. of preconditioned PKC- δ ^{+/+} hearts revealed profound changes in enzymes related to energy metabolism, e.g., NADH dehydrogenase and **ATP synthase**, with partial fragmentation of these mitochondrial enzymes and of the E2 component of the pyruvate dehydrogenase complex. Interestingly, fragmentation of mitochondrial enzymes was not observed in PKC- δ ^{-/-} hearts. High-resolution NMR anal. of cardiac metabolites demonstrated a similar rise of phosphocreatine in PKC- δ ^{+/+} and PKC- δ ^{-/-} hearts, but the preconditioning-induced increase in phosphocholine, alanine, carnitine, and glycine was restricted to PKC- δ ^{+/+} hearts, whereas lactate concns. were higher in PKC- δ ^{-/-} hearts. Taken together, the authors' results suggest that reactive oxygen species generated during ischemic preconditioning might alter mitochondrial metabolism by oxidizing key mitochondrial enzymes and that metabolic adaptation to preconditioning is impaired in PKC- δ ^{-/-} hearts.

REFERENCE COUNT: 51 THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 15 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2003:668326 CAPLUS

DOCUMENT NUMBER: 139:290251

TITLE: Ischaemic preconditioning and a mitochondrial KATP channel opener both produce cardioprotection accompanied by F1F0-ATPase inhibition in early ischaemia

AUTHOR(S): Ala-Rami, Antti; Ylitalo, Kari V.; Hassinen, Ilmo E.

CORPORATE SOURCE: Department of Medical Biochemistry and Molecular Biology, University of Oulu, Oulu, 90014, Finland

SOURCE: Basic Research in Cardiology (2003), 98(4), 250-258
CODEN: BRCAB7; ISSN: 0300-8428

PUBLISHER: Steinkopff Verlag

DOCUMENT TYPE: Journal

LANGUAGE: English

TI Ischaemic preconditioning and a mitochondrial KATP channel opener both produce cardioprotection accompanied by F1F0-ATPase inhibition in early ischaemia

AB Ischemic preconditioning gives powerful protection against prolonged ischemia affecting several intracellular regulatory and messenger pathways, although their mutual importance is far from established. Protective, preconditioning-like effects have been reported for KATP channel openers, and most of the evidence points to the mitochondrial KATP channels. We show here that the KATP channel opener diazoxide, which acts selectively on the mitochondrial channel, causes potentiation of ischemic inhibition of mitochondrial **ATP synthase** (F1F0-ATPase) along with cardioprotection. These effects are comparable with that of ischemic preconditioning. The administration of diazoxide did not affect the cellular energy state as monitored with 31P NMR. The actions of both diazoxide and ischemic preconditioning were prevented by 5-hydroxydecanoate, a specific inhibitor of the mitochondrial KATP channel. Thus mitochondrial KATP channel opening and ischemic preconditioning must share common mechanisms of action involving mitochondrial F1F0-ATPase, although involvement of the energy state in protection could not be proved.

REFERENCE COUNT: 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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ACCESSION NUMBER: 2004-0590987 PASCAL

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TITLE (IN ENGLISH): F.sub.Of.sub.1 **ATP synthase** activity is differently modulated by coronary reactive hyperemia before and after ischemic **preconditioning** in the goat

AUTHOR: PENNA Claudia; PAGLIARO Pasquale; RASTALDO Raffaella; DI PANCRAZIO Francesca; LIPPE Giovanna; GATTULLO Donatella; MANCARDI Daniele; SAMAJA Michele; LOSANO Gianni; MAVELLI Irene

CORPORATE SOURCE: Sezione di Fisiologia, Dipartimento di Neuroscienze and Dipartimento di Scienze Cliniche e Biologiche, Universita di Torino, 10100 Turin, United States; Dipartimento di Scienze e Tecnologie Biomediche and Centro di Eccellenza Micro gravity, Aging, Training, and Immobility, Universita di Udine, 33100 Udine, Italy; Dipartimento di Medicina, Chirurgia, e Odontoiatria, Universita di Milano, 20100 Milan, Italy

SOURCE: American journal of physiology. Heart and circulatory physiology, (2004), 56(5), H2192-H2200, 48 refs. ISSN: 0363-6135 CODEN: AJPPDI

DOCUMENT TYPE: Journal

BIBLIOGRAPHIC LEVEL: Analytic

COUNTRY: United States

LANGUAGE: English

AVAILABILITY: INIST-670D, 354000122491250390

TIEN F.sub.Of.sub.1 **ATP synthase** activity is differently modulated by coronary reactive hyperemia before and after ischemic **preconditioning** in the goat

AN 2004-0590987 PASCAL

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AB The amplitude of coronary reactive hyperemia (CRH), elicited by 15 s of ischemia, is reduced in hearts subjected to 5 min of ischemic **preconditioning** (IP). F.sub.Of.sub.1ATP synthase activity and ATP concentration are also altered by IP. We hypothesized that F.sub.Of.sub.1 **ATP synthase** is differently modulated by the inhibitor protein IF.sub.1 during CRH elicited before (CRH.sub.n.sub.p) and after (CRH.sub.p.sub.r.sub.e.sub.c) IP. Hemodynamic parameters were recorded in 10 anesthetized goats. Myocardial biopsies were obtained before IP (C.sub.n.sub.p), during CRH.sub.n.sub.p, 4 and 6 min after the onset of CRH.sub.n.sub.p, after IP (C.sub.p.sub.r.sub.e.sub.c), during CRH.sub.p.sub.r.sub.e.sub.c, and 4 min after CRH.sub.p.sub.r.sub.e.sub.c. F.sub.Of.sub.1ATP synthase activity, ATP concentration, and ATP-to-ADP ratio (ATP/ADP) were determined. Compared with CRH.sub.n.sub.p, IP blunted CRH.sub.p.sub.r.sub.e.sub.c. F.sub.Of.sub.1 **ATP synthase** activity transiently increased during CRH.sub.n.sub.p, decreased 4 min after CRH.sub.n.sub.p, and returned to control 2 min later; it was lower after IP (C.sub.p.sub.r.sub.e.sub.c) and did not change during and after CRH.sub.p.sub.r.sub.e.sub.c. All these changes in activity were modulated by IF.sub.1. During CRH.sub.n.sub.p, ATP concentration and ATP/ADP were reduced compared with C.sub.n.sub.p and began to rise 6 min thereafter. During C.sub.p.sub.r.sub.e.sub.c, both parameters were transiently reduced but increased during and after CRH.sub.p.sub.r.sub.e.sub.c. Hence, during CRH.sub.n.sub.p, F.sub.Of.sub.1 **ATP synthase** activity transiently increases and then decreases significantly. The shortlasting inhibition of the enzyme may explain why a few seconds of occlusion do not induce IP. After IP, F.sub.Of.sub.1 **ATP synthase** activity is blunted, and it is not affected by a subsequent 15 s of occlusion, which induces a blunted CRH.sub.p.sub.r.sub.e.sub.c. These results suggest that postischemic long-lasting inhibition of F.sub.Of.sub.1 **ATP synthase** activity may be a feature of the preconditioned heart. The increase in ATP concentration after **preconditioning** is in agreement with previous reports of reduced ATP hydrolysis by cytoplasmic

ATPases.

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ACCESSION NUMBER: 2004-0123853 PASCAL
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TITLE (IN ENGLISH): ATP-sensitive K.sup.+ channels in renal **mitochondria**
AUTHOR: CANCHERINI Douglas V.; TRABUCO Leonardo G.; REBOUCAS Nancy A.; KOWALTOWSKI Alicia J.
CORPORATE SOURCE: Departamento de Fisiologia e Biofisica, Instituto de Ciencias Biomedicas, Universidade de Sao Paulo, 05508-900 Sao Paulo, Brazil; Departamento de Bioquimica, Instituto de Quimica, Universidade de Sao Paulo, 05508-900 Sao Paulo, Brazil
SOURCE: American journal of physiology. Renal physiology, (2003), 54(6), F1291-F1296, 38 refs.
DOCUMENT TYPE: Journal
BIBLIOGRAPHIC LEVEL: Analytic
COUNTRY: United States
LANGUAGE: English
AVAILABILITY: INIST-670F, 354000118779340270

TIEN ATP-sensitive K.sup.+ channels in renal **mitochondria**
AN 2004-0123853 PASCAL
CP Copyright .COPYRGT. 2004 INIST-CNRS. All rights reserved.
AB Isolated kidney **mitochondria** swell when incubated in hyposmotic solutions containing K.sup.+ salts in a manner inhibited by ATP, ADP, 5-hydroxydecanoate, and glibenclamide and stimulated by GTP and diazoxide. These results suggest the existence of ATP-sensitive K.sup.+ channels in these **mitochondria**, similar to those previously described in heart, liver, and brain. Renal mitochondrial ATP-sensitive K.sup.+ uptake rates are .eqvsim.140 nmol.min.sup.-.sup.1.mg protein.sup.-.sup.1. This K.sup.+ transport results in a slight increase in respiration and decrease in the inner membrane potential. In addition, the activation of ATP-inhibited K.sup.+ uptake using diazoxide leads to a decrease of ATP hydrolysis through the reverse activity of the F.sub.0F.sub.1 **ATP synthase** when respiration is inhibited. In conclusion, we characterize an ATP-sensitive K.sup.+ transport pathway in kidney **mitochondria** that affects volume, respiration, and membrane potential and may have a role in the prevention of mitochondrial ATP hydrolysis.

L4 ANSWER 8 OF 15 USPATFULL on STN

ACCESSION NUMBER: 2004:327968 USPATFULL
TITLE: Methods and compositions for modulating proteins modified in preconditioning against ischemia/hypoxia
INVENTOR(S): Eyk, Jennifer E. Van, Baltimore, MD, UNITED STATES
Elliott, Steven T., Cockeysville, MD, UNITED STATES
Arrell, David Kent, Rochester, MN, UNITED STATES

	NUMBER	KIND	DATE
PATENT INFORMATION:	US 2004259793	A1	20041223
APPLICATION INFO.:	US 2004-824027	A1	20040414 (10)

	NUMBER	DATE
PRIORITY INFORMATION:	US 2003-463139P	20030414 (60)
DOCUMENT TYPE:	Utility	
FILE SEGMENT:	APPLICATION	
LEGAL REPRESENTATIVE:	Licata & Tyrrell P.C., 66 East Main Street, Marlton, NJ, 08053	

NUMBER OF CLAIMS: 12
EXEMPLARY CLAIM: 1
NUMBER OF DRAWINGS: 12 Drawing Page(s)
LINE COUNT: 1334

TI Methods and compositions for modulating proteins modified in preconditioning against ischemia/hypoxia
AB Proteins modified by pharmacological preconditioning are provided. Compositions, methods and events for modulating these proteins and priming cells for preconditioning and inducing preconditioning in a cell, tissue or organ as well as methods for identifying new compositions and methods for such priming and induction are also provided. In addition, methods for diagnosing and monitoring preconditioning or ischemic, hypoxic, ischemic/reperfusion and hypoxic/reperfusion conditions or the ability of a cell, tissue or organ to survive injury by measuring modulation of one or more of these preconditioning proteins are provided.

L4 ANSWER 9 OF 15 USPATFULL on STN

ACCESSION NUMBER: 2003:176318 USPATFULL
TITLE: Methods to identify compounds affecting mitochondria
INVENTOR(S): Marban, Eduardo, Lutherville, MD, United States
O'Rourke, Brian, Sparks, MD, United States
PATENT ASSIGNEE(S): Johns Hopkins University, Baltimore, MD, United States
(U.S. corporation)

	NUMBER	KIND	DATE
PATENT INFORMATION:	US 6586241	B1	20030701
APPLICATION INFO.:	US 2000-684730		20001006 (9)
RELATED APPLN. INFO.:	Continuation of Ser. No. US 1998-60774, filed on 15 Apr 1998, now patented, Pat. No. US 6183948		
DOCUMENT TYPE:	Utility		
FILE SEGMENT:	GRANTED		
PRIMARY EXAMINER:	Lankford, Jr., Leon B.		
LEGAL REPRESENTATIVE:	Corless, Peter F., Edwards & Angell, LLP		
NUMBER OF CLAIMS:	3		
EXEMPLARY CLAIM:	1		
NUMBER OF DRAWINGS:	14 Drawing Figure(s); 6 Drawing Page(s)		
LINE COUNT:	1039		

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

TI Methods to identify compounds affecting mitochondria
AB The present invention relates to methods for identifying a compound capable of modulating mitochondrial function, comprising contacting a eukaryotic cell with one or more candidate compounds, and detecting a change in the mitochondrial redox state of the cell. The methods further relates to such methods wherein endogenous fluorescence of the cell mitochondria is indicative of a change of redox state.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L4 ANSWER 10 OF 15 USPATFULL on STN

ACCESSION NUMBER: 2003:30273 USPATFULL
TITLE: Methods and compositions for modulating adenosine triphosphate (ATP) in cells and preventing cell injury or death via post-translational modifications to **ATP synthase**
INVENTOR(S): Van Eyk, Jennifer E., Kingston, CANADA
Arrell, David Kent, Kingston, CANADA

	NUMBER	KIND	DATE
PATENT INFORMATION:	US 2003022220	A1	20030130

APPLICATION INFO.: US 2002-189820 A1 20020703 (10)

	NUMBER	DATE
PRIORITY INFORMATION:	US 2001-303491P	20010706 (60)
DOCUMENT TYPE:	Utility	
FILE SEGMENT:	APPLICATION	
LEGAL REPRESENTATIVE:	Licata & Tyrrell P.C., 66 E. Main Street, Marlton, NJ, 08053	
NUMBER OF CLAIMS:	4	
EXEMPLARY CLAIM:	1	
NUMBER OF DRAWINGS:	8 Drawing Page(s)	
LINE COUNT:	762	

opp and 112(e)

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

TI Methods and compositions for modulating adenosine triphosphate (ATP) in cells and preventing cell injury or death via post-translational modifications to **ATP synthase**

AB Compositions and methods for modulating adenosine triphosphate (ATP) in cells via altering post-translational modifications of **ATP synthase** subunits or precursors thereof such as the **ATP synthase β** chain and its precursor are provided. These compositions and methods are useful in preconditioning organs and preventing cell injury or cell death via regulating ATP synthesis or hydrolysis in cells of the organs.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L4 ANSWER 11 OF 15 USPATFULL on STN

ACCESSION NUMBER: 2001:18196 USPATFULL

TITLE: Methods to identify compounds affecting mitochondria

INVENTOR(S): Marban, Eduardo, Lutherville, MD, United States
O'Rourke, Brian, Sparks, MD, United States

PATENT ASSIGNEE(S): Johns Hopkins University, Baltimore, MD, United States
(U.S. corporation)

	NUMBER	KIND	DATE
PATENT INFORMATION:	US 6183948	B1	20010206
APPLICATION INFO.:	US 1998-60774		19980415 (9)
DOCUMENT TYPE:	Utility		
FILE SEGMENT:	Granted		
PRIMARY EXAMINER:	Lankford, Jr., Leon B.		
LEGAL REPRESENTATIVE:	Corless, Peter F., Schray, Kerri P. Edwards & Angell, LLP.		
NUMBER OF CLAIMS:	31		
EXEMPLARY CLAIM:	1		
NUMBER OF DRAWINGS:	14 Drawing Figure(s); 6 Drawing Page(s)		
LINE COUNT:	1117		

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

TI Methods to identify compounds affecting mitochondria

AB The present invention relates to methods for identifying a compound capable of modulating mitochondrial function, comprising contacting a eukaryotic cell with one or more candidate compounds, and detecting a change in the mitochondrial redox state of the cell. The methods further relates to such methods wherein endogenous fluorescence of the cell mitochondria is indicative of a change of redox state.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L4 ANSWER 12 OF 15 Elsevier BIOBASE COPYRIGHT 2005 Elsevier Science B.V. on STN

ACCESSION NUMBER: 2004278351 ESBIODASE

TITLE: F.sub.0F.sub.1 **ATP synthase**

activity is differently modulated by coronary reactive hyperemia before and after ischemic **preconditioning** in the goat

AUTHOR: Penna C.; Pagliaro P.; Rastaldo R.; Di Pancrazio F.; Lippe G.; Gattullo D.; Mancardi D.; Samaja M.; Losano G.; Mavelli I.

CORPORATE SOURCE: P. Pagliaro, Dipto. di Sci. Cliniche e Biologiche, Universita di Torino, Ospedale S. Luigi, Regione Gonzole, 10043 Orbassano (TO), Italy.
E-mail: pasquale.pagliaro@unito.it

SOURCE: American Journal of Physiology - Heart and Circulatory Physiology, (2004), 287/5 56-5 (H2192-H2200), 48 reference(s)
CODEN: AJPPDI ISSN: 0363-6135

DOCUMENT TYPE: Journal; Article

COUNTRY: United States

LANGUAGE: English

SUMMARY LANGUAGE: English

TI F.sub.0F.sub.1 **ATP synthase** activity is differently modulated by coronary reactive hyperemia before and after ischemic **preconditioning** in the goat

AB The amplitude of coronary reactive hyperemia (CRH), elicited by 15 s of ischemia, is reduced in hearts subjected to 5 min of ischemic **preconditioning** (IP). F.sub.0F.sub.1 **ATP synthase** activity and ATP concentration are also altered by IP. We hypothesized that F.sub.0F.sub.1 **ATP synthase** is differently modulated by the inhibitor protein IF.sub.1 during CRH elicited before (CRH.sub.n.sub.p) and after (CRH.sub.p.sub.r.sub.e.sub.c) IP. Hemodynamic parameters were recorded in 10 anesthetized goats. Myocardial biopsies were obtained before IP (C.sub.n.sub.p), during CRH.sub.n.sub.p, 4 and 6 min after the onset of CRH.sub.n.sub.p, after IP (C.sub.p.sub.r.sub.e.sub.c), during CRH.sub.p.sub.r.sub.e.sub.c, and 4 min after CRH.sub.p.sub.r.sub.e.sub.c. F.sub.0F.sub.1 **ATP synthase** activity, ATP concentration, and ATP-to-ADP ratio (ATP/ADP) were determined. Compared with CRH.sub.n.sub.p, IP blunted CRH.sub.p.sub.r.sub.e.sub.c. F.sub.0F.sub.1 **ATP synthase** activity transiently increased during CRH.sub.n.sub.p, decreased 4 min after CRH .sub.n.sub.p, and returned to control 2 min later; it was lower after IP (C.sub.p.sub.r.sub.e.sub.c) and did not change during and after CRH.sub.p.sub.r.sub.e.sub.c. All these changes in activity were modulated by IF.sub.1. During CRH .sub.n.sub.p, ATP concentration and ATP/ADP were reduced compared with C .sub.n.sub.p and began to rise 6 min thereafter. During C.sub.p.sub.r.sub.e.sub.c, both parameters were transiently reduced but increased during and after CRH .sub.p.sub.r.sub.e.sub.c. Hence, during CRH.sub.n.sub.p, F.sub.0F.sub.1 **ATP synthase** activity transiently increases and then decreases significantly. The short-lasting inhibition of the enzyme may explain why a few seconds of occlusion do not induce IP. After IP, F.sub.0F.sub.1 **ATP synthase** activity is blunted, and it is not affected by a subsequent 15 s of occlusion, which induces a blunted CRH.sub.p.sub.r.sub.e.sub.c. These results suggest that postischemic long-lasting inhibition of F.sub.0F.sub.1 **ATP synthase** activity may be a feature of the preconditioned heart. The increase in ATP concentration after **preconditioning** is in agreement with previous reports of reduced ATP hydrolysis by cytoplasmic ATPases.

L4 ANSWER 13 OF 15 Elsevier BIOBASE COPYRIGHT 2005 Elsevier Science B.V. on STN

ACCESSION NUMBER: 2004213219 ESBIOBASE

TITLE: Multiprotein complex containing succinate dehydrogenase confers mitochondrial ATP-sensitive K.sup.+ channel activity

AUTHOR: Ardehali H.; Chen Z.; Ko Y.; Mejia-Alvarez R.; Marban E.
CORPORATE SOURCE: E. Marban, 844 Ross Building, Johns Hopkins University, 720 Rutland Avenue, Baltimore, MD 21205, United States.
E-mail: marban@jhmi.edu
SOURCE: Proceedings of the National Academy of Sciences of the United States of America, (10 AUG 2004), 101/32 (11880-11885), 31 reference(s)
CODEN: PNASA6 ISSN: 0027-8424
DOCUMENT TYPE: Journal; Article
COUNTRY: United States
LANGUAGE: English
SUMMARY LANGUAGE: English

TI Multiprotein complex containing succinate dehydrogenase confers mitochondrial ATP-sensitive K_{sup}.+ channel activity
AB The mitochondrial ATP-sensitive K_{sup}.+ (mitoK_{sub}.A_{sub}.T_{sub}.P) channel plays a central role in protection of cardiac and neuronal cells against ischemia and apoptosis, but its molecular structure is unknown. Succinate dehydrogenase (SDH) is inhibited by mitoK_{sub}.A_{sub}.T_{sub}.P activators, fueling the contrary view that SDH, rather than mitoK_{sub}.A_{sub}.T_{sub}.P, is the target of cardioprotective drugs. Here, we report that SDH forms part of mitoK_{sub}.A_{sub}.T_{sub}.P functionally and structurally. Four mitochondrial proteins [mitochondrial ATP-binding cassette protein 1 (mABC1), phosphate carrier, adenine nucleotide translocator, and **ATP synthase**] associate with SDH. A purified IM fraction containing these proteins was reconstituted into proteoliposomes and lipid bilayers and shown to confer mitoK_{sub}.A_{sub}.T_{sub}.P channel activity. This channel activity is sensitive not only to mitoK_{sub}.A_{sub}.T_{sub}.P activators and blockers but also to SDH inhibitors. These results reconcile the controversy over the basis of ischemic **preconditioning** by demonstrating that SDH is a component of mitoK_{sub}.A_{sub}.T_{sub}.P as part of a macromolecular super-complex. The findings also provide a tangible clue as to the structural basis of mitoK_{sub}.A_{sub}.T_{sub}.P channels.

L4 ANSWER 14 OF 15 Elsevier BIOBASE COPYRIGHT 2005 Elsevier Science B.V. on STN

ACCESSION NUMBER: 2003289067 ESBIODASE
TITLE: ATP-sensitive K_{sup}.+ channels in renal **mitochondria**
AUTHOR: Cancherini D.V.; Trabuco L.G.; Reboucas N.A.; Kowaltowski A.J.
CORPORATE SOURCE: A.J. Kowaltowski, Cidade Universitaria, Av. Prof. Lineu Prestes, 748, 05508-900, Sao Paulo, Brazil.
E-mail: alicia@iq.usp.br
SOURCE: American Journal of Physiology - Renal Physiology, (2003), 285/6 54-6 (F1291-F1296), 38 reference(s)
CODEN: AJPPFK ISSN: 0363-6127
DOCUMENT TYPE: Journal; Article
COUNTRY: United States
LANGUAGE: English
SUMMARY LANGUAGE: English
TI ATP-sensitive K_{sup}.+ channels in renal **mitochondria**
AB Isolated kidney **mitochondria** swell when incubated in hyposmotic solutions containing K_{sup}.+ salts in a manner inhibited by ATP, ADP, 5-hydroxydecanoate, and glibenclamide and stimulated by GTP and diazoxide. These results suggest the existence of ATP-sensitive K_{sup}.+ channels in these **mitochondria**, similar to those previously described in heart, liver, and brain. Renal mitochondrial ATP-sensitive K_{sup}.+ uptake rates are .apprx.140 nmol.midldot.min.sup.-.sup.1-mg protein.sup.-.sup.1. This K_{sup}.+ transport results in a slight increase in respiration and decrease in the inner membrane potential. In addition,

the activation of ATP-inhibited K^{sup.}+ uptake using diazoxide leads to a decrease of ATP hydrolysis through the reverse activity of the F₁sub.0F₁sub.1 **ATP synthase** when respiration is inhibited. In conclusion, we characterize an ATP-sensitive K^{sup.}+ transport pathway in kidney **mitochondria** that affects volume, respiration, and membrane potential and may have a role in the prevention of mitochondrial ATP hydrolysis.

L4 ANSWER 15 OF 15 Elsevier BIOBASE COPYRIGHT 2005 Elsevier Science B.V.
on STN

ACCESSION NUMBER: 2002273905 ESBIOBASE
TITLE: Halothane, isoflurane and sevoflurane inhibit **NADH: Ubiquinone oxidoreductase** (complex I) of cardiac **mitochondria**
AUTHOR: Hanley P.J.; Ray J.; Brandt U.; Daut J.
CORPORATE SOURCE: J. Daut, Institute of Physiology, Marburg University, Deutschhausstrasse 2, 35037 Marburg, Germany.
E-mail: daut@mail.uni-marburg.de
SOURCE: Journal of Physiology, (01 NOV 2002), 544/3 (687-693), 40 reference(s)
CODEN: JPHYA7 ISSN: 0022-3751
DOCUMENT TYPE: Journal; Article
COUNTRY: United Kingdom
LANGUAGE: English
SUMMARY LANGUAGE: English

TI Halothane, isoflurane and sevoflurane inhibit **NADH: Ubiquinone oxidoreductase** (complex I) of cardiac **mitochondria**

AB We have investigated the effects of volatile anaesthetics on electron transport chain activity in the mammalian heart. Halothane, isoflurane and sevoflurane reversibly increased NADH fluorescence (autofluorescence) in intact ventricular myocytes of guinea-pig, suggesting that NADH oxidation was impaired. Using pig heart submitochondrial particles we found that the anaesthetics dose-dependently inhibited NADH oxidation in the order: halothane > isoflurane = sevoflurane. Succinate oxidation was unaffected by either isoflurane or sevoflurane, indicating that these agents selectively inhibit complex I (**NADH:ubiquinone oxidoreductase**). In addition to inhibiting NADH oxidation, halothane also inhibited succinate oxidation (and succinate dehydrogenase), albeit to a lesser extent. To test the hypothesis that complex I is a target of volatile anaesthetics, we examined the effects of these agents on **NADH:ubiquinone oxidoreductase** (EC 1.6.99.3) activity using the ubiquinone analogue DBQ (decylubiquinone) as substrate. Halothane, isoflurane and sevoflurane dose-dependently inhibited NADH:DBQ oxidoreductase activity. Unlike the classical inhibitor rotenone, none of the anaesthetics completely inhibited enzyme activity at high concentration, suggesting that these agents bind weakly to the 'hydrophobic inhibitory site' of complex I. In conclusion, halothane, isoflurane and sevoflurane inhibit complex I (**NADH:ubiquinone oxidoreductase**) of the electron transport chain. At concentrations of .apprx.2 MAC (minimal alveolar concentration), the activity of **NADH: ubiquinone oxidoreductase** was reduced by about 20% in the presence of halothane or isoflurane, and by about 10% in the presence of sevoflurane. These inhibitory effects are unlikely to compromise cardiac performance at usual clinical concentrations, but may contribute to the mechanism by which volatile anaesthetics induce pharmacological **preconditioning**.

=> s vaneyk,j?/au

L5 72 VANEYK,J?/AU

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=> s arrell,d?/au
L6          76 ARRELL,D?/AU

=> s elliot,s?/au
L7          3875 ELLIOTT,S?/AU

=> s L5 AND L6 AND L7
L8          0 L5 AND L6 AND L7

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L9          6 EYK,J?/AU

=> s L9 AND L7 AND L8
L10         0 L9 AND L7 AND L8

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(FILE 'HOME' ENTERED AT 14:23:43 ON 09 MAY 2005)

INDEX 'ADISCTI, ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, ANTE, AQUALINE, AQUASCI, BIOBUSINESS, BIOCOMMERCE, BIOENG, BIOSIS, BIOTECHABS, BIOTECHDS, BIOTECHNO, CABA, CANCERLIT, CAPLUS, CEABA-VTB, CEN, CIN, CONFSCI, CROPB, CROPU, DDFB, DDFU, DGENE, DISSABS, ...' ENTERED AT 14:23:54 ON 09 MAY 2005
SEA PRECONDITIONING

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370  FILE ADISCTI
18   FILE ADISINSIGHT
14   FILE ADISNEWS
275  FILE AGRICOLA
70   FILE ANABSTR
14   FILE ANTE
27   FILE AQUALINE
140  FILE AQUASCI
121  FILE BIOBUSINESS
4    FILE BIOCOMMERCE
131  FILE BIOENG
6800 FILE BIOSIS
109  FILE BIOTECHABS
109  FILE BIOTECHDS
412  FILE BIOTECHNO
657  FILE CABA
315  FILE CANCERLIT
5254 FILE CAPLUS
54   FILE CEABA-VTB
15   FILE CIN
267  FILE CONFSCI
15   FILE CROPB
59   FILE CROPU
5    FILE DDFB
869  FILE DDFU
76   FILE DGENE
553  FILE DISSABS
5    FILE DRUGB
994  FILE DRUGU
82   FILE EMBAL
4241 FILE EMBASE
2600 FILE ESBIODASE
206  FILE FEDRIP
65   FILE FROSTI
98   FILE FSTA
61   FILE GENBANK
9    FILE HEALSAFE
```



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1201  FILE IFIPAT
      3  FILE IMSDRUGNEWS
      4  FILE IMSRESEARCH
    711  FILE JICST-EPLUS
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    526  FILE LIFESCI
      6  FILE MEDICONF
   4881  FILE MEDLINE
      24  FILE NIOSHTIC
    583  FILE NTIS
      75  FILE OCEAN
   5144  FILE PASCAL
      2  FILE PHAR
      3  FILE PHARMAML
     18  FILE PHIN
    306  FILE PROMT
      6  FILE PROUSDDR
      9  FILE RDISCLOSURE
   8243  FILE SCISEARCH
   2045  FILE TOXCENTER
   4450  FILE USPATFULL
     361  FILE USPAT2
      8  FILE VETB
     14  FILE VETU
     99  FILE WATER
    718  FILE WPIDS
      3  FILE WPIFV
    718  FILE WPINDEX

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L1 QUE PRECONDITIONING

FILE 'SCISEARCH, BIOSIS, CAPLUS, PASCAL, MEDLINE, USPATFULL, EMBASE,
ESBIOBASE, TOXCENTER' ENTERED AT 14:25:03 ON 09 MAY 2005

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L3      508 DUP REM L2 (753 DUPLICATES REMOVED)
L4      15 S L3 AND (ISOCITRATE(W) DEHYDROGENASE (W) NAD (W) ALPHA (W) SUBUNIT O
L5      72 S VANEYK,J?/AU
L6      76 S ARRELL,D?/AU
L7      3875 S ELLIOTT,S?/AU
L8      0 S L5 AND L6 AND L7
L9      6 S EYK,J?/AU
L10     0 S L9 AND L7 AND L8

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L13  ANSWER 1 OF 2      MEDLINE on STN
TI   [Molecular modifications of troponin I and T detected in serum from
      patients with acute myocardial infarction].
      Molekylaere forandringer af troponin I og T pavist i serum hos patienter
      med akut myokardieinfarkt.

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L13  ANSWER 2 OF 2      BIOSIS  COPYRIGHT (c) 2005 The Thomson Corporation  on STN
TI   Identification of myofilament proteins that are degraded and/or released

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during ischemia/reperfusion: Effects on the pCa-force relation.

=> d l13 ibib ti abs 1-2

L13 ANSWER 1 OF 2 MEDLINE on STN
ACCESSION NUMBER: 2003044477 MEDLINE
DOCUMENT NUMBER: PubMed ID: 12553091
TITLE: [Molecular modifications of troponin I and T detected in serum from patients with acute myocardial infarction]. Molekylaere forandringer af troponin I og T pavist i serum hos patienter med akut myokardieinfarkt.
AUTHOR: Atar Dan; Madsen Lene Helleskov; Labugger Ralf; **VanEyck Jennifer E**
CORPORATE SOURCE: H:S Frederiksberg Hospital, kardiologisk klinik E, DK-2000 Frederiksberg.. datar@dadlnet.dk
SOURCE: Ugeskrift for laeger, (2003 Jan 6) 165 (2) 116-20. Journal code: 0141730. ISSN: 0041-5782.
PUB. COUNTRY: Denmark
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)
LANGUAGE: Danish
FILE SEGMENT: Priority Journals
ENTRY MONTH: 200302
ENTRY DATE: Entered STN: 20030130
Last Updated on STN: 20030228
Entered Medline: 20030227
TI [Molecular modifications of troponin I and T detected in serum from patients with acute myocardial infarction]. Molekylaere forandringer af troponin I og T pavist i serum hos patienter med akut myokardieinfarkt.
AB INTRODUCTION: Cardiac troponin I and T (cTnI and cTnT) are specific biochemical serum markers for acute myocardial infarction (AMI). However, cTnI diagnostic assays are plagued by difficulties, resulting in > 20-fold differences in measured values. These discrepancies may result from the release of the numerous cTnI modification products that are present in ischaemic myocardium. The resolution of these discrepancies requires an investigation of the exact forms of the troponins present in the bloodstream of patients after myocardial injury. MATERIAL AND METHODS: A Westernblot direct serum analysis protocol was developed that allowed us to detect intact cTnI and a spectrum of up to 11 modified products in the serum from patients with AMI. RESULTS: We document both a cTnI degradation pattern and the existence of phosphorylated cTnI in serum. The number and extent of these modifications reflect patterns similar to the time profiles of the routine clinical serum markers of total creatine kinase, creatine kinase-MB, and cTnI (determined by ELISA). Data from in vitro experiments, which were undertaken to study the degradation of human recombinant cTnI and cTnT when spiked in serum, indicate that some modification products present in patient serum existed in the myocardium. DISCUSSION: This pilot study defines, for the first time, what forms of cTnI and cTnT appear in the bloodstream of AMI patients, and it clarifies the lack of standardization between different cTnI diagnostic assays.

L13 ANSWER 2 OF 2 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN
ACCESSION NUMBER: 1997:4179 BIOSIS
DOCUMENT NUMBER: PREV199799303382
TITLE: Identification of myofilament proteins that are degraded and/or released during ischemia/reperfusion: Effects on the pCa-force relation.
AUTHOR(S): **Vaneyk, Jennifer E.** [Reprint author]; Powers, Francis M.; Law, William R.; Hodges, Robert S.; Solaro, R. John
CORPORATE SOURCE: Univ. Ill., Chicago, IL, USA
SOURCE: Circulation, (1996) Vol. 94, No. 8 SUPPL., pp. I365.

Meeting Info.: 69th Scientific Sessions of the American
Heart Association. New Orleans, Louisiana, USA. November
10-13, 1996.

CODEN: CIRCAZ. ISSN: 0009-7322.

DOCUMENT TYPE:

Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE:

English

ENTRY DATE:

Entered STN: 7 Jan 1997

Last Updated on STN: 7 Jan 1997

TI Identification of myofilament proteins that are degraded and/or released
during ischemia/reperfusion: Effects on the pCa-force relation.

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(FILE 'HOME' ENTERED AT 14:23:43 ON 09 MAY 2005)

INDEX 'ADISCTI, ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, ANTE, AQUALINE,
AQUASCI, BIOBUSINESS, BIOCOMMERCE, BIOENG, BIOSIS, BIOTECHABS, BIOTECHDS,
BIOTECHNO, CABA, CANCERLIT, CAPLUS, CEABA-VTB, CEN, CIN, CONFSCI, CROPB,
CROPU, DDFB, DDFU, DGENE, DISSABS, ...' ENTERED AT 14:23:54 ON 09 MAY 2005
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14 FILE ANTE
27 FILE AQUALINE
140 FILE AQUASCI
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4 FILE BIOCOMMERCE
131 FILE BIOENG
6800 FILE BIOSIS
109 FILE BIOTECHABS
109 FILE BIOTECHDS
412 FILE BIOTECHNO
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5254 FILE CAPLUS
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15 FILE CIN
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15 FILE CROPB
59 FILE CROPU
5 FILE DDFB
869 FILE DDFU
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5 FILE DRUGB
994 FILE DRUGU
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4241 FILE EMBASE
2600 FILE ESBIODASE
206 FILE FEDRIP
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9 FILE HEALSAFE
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4 FILE IMSRESEARCH

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6 FILE PROUSDDR
9 FILE RDISCLOSURE
8243 FILE SCISEARCH
2045 FILE TOXCENTER
4450 FILE USPATFULL
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99 FILE WATER
718 FILE WPIDS
3 FILE WPIFV
718 FILE WPINDEX
L1 QUE PRECONDITIONING
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FILE 'SCISEARCH, BIOSIS, CAPLUS, PASCAL, MEDLINE, USPATFULL, EMBASE,
ESBIOBASE, TOXCENTER' ENTERED AT 14:25:03 ON 09 MAY 2005
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L3 508 DUP REM L2 (753 DUPLICATES REMOVED)
L4 15 S L3 AND (ISOCITRATE(W) DEHYDROGENASE(W) NAD(W) ALPHA(W) SUBUNIT O
L5 72 S VANEYK,J?/AU
L6 76 S ARRELL,D?/AU
L7 3875 S ELLIOTT,S?/AU
L8 0 S L5 AND L6 AND L7
L9 6 S EYK,J?/AU
L10 0 S L9 AND L7 AND L8
L11 0 S VANEYK,JENNIFER/AU
L12 2 S VANEYK,JENNIFER?/AU
L13 2 DUP REM L12 (0 DUPLICATES REMOVED)

```

```

=> s L6 AND L7
L14 5 L6 AND L7

```

```

=> dup rem l14
PROCESSING COMPLETED FOR L14
L15 5 DUP REM L14 (0 DUPLICATES REMOVED)

```

```

=> d l15 ibib ti abs 1-5

```

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L15 ANSWER 1 OF 5 USPATFULL on STN
ACCESSION NUMBER: 2004:327968 USPATFULL
TITLE: Methods and compositions for modulating proteins
modified in preconditioning against ischemia/hypoxia
INVENTOR(S): Eyk, Jennifer E. Van, Baltimore, MD, UNITED STATES
Elliot, Steven T., Cockeysville, MD, UNITED STATES
Arrell, David Kent, Rochester, MN, UNITED STATES

```

```

NUMBER KIND DATE

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PATENT INFORMATION: US 2004259793 A1 20041223
APPLICATION INFO.: US 2004-824027 A1 20040414 (10)

	NUMBER	DATE
PRIORITY INFORMATION:	US 2003-463139P	20030414 (60)
DOCUMENT TYPE:	Utility	
FILE SEGMENT:	APPLICATION	
LEGAL REPRESENTATIVE:	Licata & Tyrrell P.C., 66 East Main Street, Marlton, NJ, 08053	
NUMBER OF CLAIMS:	12	
EXEMPLARY CLAIM:	1	
NUMBER OF DRAWINGS:	12 Drawing Page(s)	
LINE COUNT:	1334	
TI	Methods and compositions for modulating proteins modified in preconditioning against ischemia/hypoxia	
AB	Proteins modified by pharmacological preconditioning are provided. Compositions, methods and events for modulating these proteins and priming cells for preconditioning and inducing preconditioning in a cell, tissue or organ as well as methods for identifying new compositions and methods for such priming and induction are also provided. In addition, methods for diagnosing and monitoring preconditioning or ischemic, hypoxic, ischemic/reperfusion and hypoxic/reperfusion conditions or the ability of a cell, tissue or organ to survive injury by measuring modulation of one or more of these preconditioning proteins are provided.	

L15 ANSWER 2 OF 5 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN
ACCESSION NUMBER: 2004:40526 BIOSIS
DOCUMENT NUMBER: PREV200400041897
TITLE: 2-Dimensional gel electrophoresis proteomic database of rabbit ventricular myocytes.

AUTHOR(S): **Elliott, S.** [Reprint Author]; **Arrell, D. Kent** [Reprint Author]; Doherty-Kirby, A.; Brown, H. [Reprint Author]; Lajoie, G.; Marban, E.; Van Eyk, J. [Reprint Author]

CORPORATE SOURCE: Queen's University, Kingston, ON, Canada
SOURCE: Molecular & Cellular Proteomics, (September 2003) Vol. 2, No. 9, pp. 835. print.
Meeting Info.: HUPO (Human Proteomics Organisation) 2nd Annual and IUBMB (International Union of Biochemistry and Molecular Biology) XIX World Congress. Montreal, Quebec, Canada. October 08-11, 2003. American Society for Biochemistry and Molecular Biology Inc.
ISSN: 1535-9476 (ISSN print).

DOCUMENT TYPE: Conference; (Meeting)
Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 14 Jan 2004
Last Updated on STN: 14 Jan 2004

TI 2-Dimensional gel electrophoresis proteomic database of rabbit ventricular myocytes.

L15 ANSWER 3 OF 5 SCISEARCH COPYRIGHT (c) 2005 The Thomson Corporation on STN

ACCESSION NUMBER: 2002:162755 SCISEARCH

THE GENUINE ARTICLE: 511EY

TITLE: Preconditioning post-translationally modifies cardiac mitochondrial F1Fo ATPase subunit

AUTHOR: **Arrell D K (Reprint); Neverova I; Elliott S T; Turcotte A; Van Eyk J E**

CORPORATE SOURCE: Queens Univ, Kingston, ON K7L 3N6, Canada
 COUNTRY OF AUTHOR: Canada
 SOURCE: BIOPHYSICAL JOURNAL, (JAN 2002) Vol. 82, No. 1, Part 2, pp. 611A-611A. MA 2986.
 Publisher: BIOPHYSICAL SOCIETY, 9650 ROCKVILLE PIKE, BETHESDA, MD 20814-3998 USA.
 ISSN: 0006-3495.

DOCUMENT TYPE: Conference; Journal
 LANGUAGE: English
 REFERENCE COUNT: 0

TI Preconditioning post-translationally modifies cardiac mitochondrial F1Fo ATPase subunit

L15 ANSWER 4 OF 5 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN
 ACCESSION NUMBER: 2002:365381 BIOSIS
 DOCUMENT NUMBER: PREV200200365381
 TITLE: Preconditioning post-translationally modifies cardiac mitochondrial F1Fo ATPase beta subunit.

AUTHOR(S): **Arrell, D. Kent** [Reprint author]; Neverova, Irina [Reprint author]; **Elliott, Steven T.** [Reprint author]; Turcotte, Antony [Reprint author]; Van Eyk, Jennifer E. [Reprint author]

CORPORATE SOURCE: Queen's University, 414 Botterell Hall, Kingston, Ontario, K7L 3N6, Canada

SOURCE: Biophysical Journal, (January, 2002) Vol. 82, No. 1 Part 2, pp. 611a. print.
 Meeting Info.: 46th Annual Meeting of the Biophysical Society. San Francisco, California, USA. February 23-27, 2002.
 CODEN: BIOJAU. ISSN: 0006-3495.

DOCUMENT TYPE: Conference; (Meeting)
 Conference; Abstract; (Meeting Abstract)
 Conference; (Meeting Poster)

LANGUAGE: English
 ENTRY DATE: Entered STN: 3 Jul 2002
 Last Updated on STN: 3 Jul 2002

TI Preconditioning post-translationally modifies cardiac mitochondrial F1Fo ATPase beta subunit.

L15 ANSWER 5 OF 5 SCISEARCH COPYRIGHT (c) 2005 The Thomson Corporation on STN
 ACCESSION NUMBER: 2002:687876 SCISEARCH
 THE GENUINE ARTICLE: 579YR
 TITLE: Proteomic analysis of preconditioning reveals post-translational modification of the F1Fo ATPase beta subunit

AUTHOR: **Arrell D K (Reprint); Elliott S;**
 Neverova I; Marban E; Van Eyk J E

CORPORATE SOURCE: Queens Univ, Dept Physiol, Kingston, ON, Canada; Queens Univ, Dept Biochem, Kingston, ON, Canada; Johns Hopkins Univ, Inst Mol Cardiobiol, Baltimore, MD USA

COUNTRY OF AUTHOR: Canada; USA

SOURCE: JOURNAL OF MOLECULAR AND CELLULAR CARDIOLOGY, (JUL 2002) Vol. 34, No. 7, pp. A33-A33.
 Publisher: ACADEMIC PRESS LTD ELSEVIER SCIENCE LTD, 24-28 OVAL RD, LONDON NW1 7DX, ENGLAND.
 ISSN: 0022-2828.

DOCUMENT TYPE: Conference; Journal
 LANGUAGE: English
 REFERENCE COUNT: 0

TI Proteomic analysis of preconditioning reveals post-translational modification of the F1Fo ATPase beta subunit

=> FIL STNGUIDE

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

132.69

134.08

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE

TOTAL

ENTRY

SESSION

CA SUBSCRIBER PRICE

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-1.46

FILE 'STNGUIDE' ENTERED AT 14:38:33 ON 09 MAY 2005

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AND TECHNOLOGY CORPORATION, AND FACHINFORMATIONSZENTRUM KARLSRUHE

FILE CONTAINS CURRENT INFORMATION.

LAST RELOADED: May 6, 2005 (20050506/UP).

=> d his

(FILE 'HOME' ENTERED AT 14:23:43 ON 09 MAY 2005)

INDEX 'ADISCTI, ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, ANTE, AQUALINE, AQUASCI, BIOBUSINESS, BIOCOMMERCE, BIOENG, BIOSIS, BIOTECHABS, BIOTECHDS, BIOTECHNO, CABA, CANCERLIT, CAPLUS, CEABA-VTB, CEN, CIN, CONFSCI, CROPB, CROPU, DDFB, DDFU, DGENE, DISSABS, ...' ENTERED AT 14:23:54 ON 09 MAY 2005

SEA PRECONDITIONING

370 FILE ADISCTI
18 FILE ADISINSIGHT
14 FILE ADISNEWS
275 FILE AGRICOLA
70 FILE ANABSTR
14 FILE ANTE
27 FILE AQUALINE
140 FILE AQUASCI
121 FILE BIOBUSINESS
4 FILE BIOCOMMERCE
131 FILE BIOENG
6800 FILE BIOSIS
109 FILE BIOTECHABS
109 FILE BIOTECHDS
412 FILE BIOTECHNO
657 FILE CABA
315 FILE CANCERLIT
5254 FILE CAPLUS
54 FILE CEABA-VTB
15 FILE CIN
267 FILE CONFSCI
15 FILE CROPB
59 FILE CROPU
5 FILE DDFB
869 FILE DDFU
76 FILE DGENE
553 FILE DISSABS
5 FILE DRUGB
994 FILE DRUGU
82 FILE EMBAL
4241 FILE EMBASE
2600 FILE ESBIODASE
206 FILE FEDRIP
65 FILE FROSTI
98 FILE FSTA

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        61  FILE GENBANK
          9  FILE HEALSAFE
1201      FILE IFIPAT
          3  FILE IMSDRUGNEWS
          4  FILE IMSRESEARCH
       711  FILE JICST-EPLUS
          7  FILE KOSMET
       526  FILE LIFESCI
          6  FILE MEDICONF
     4881  FILE MEDLINE
          24  FILE NIOSHTIC
       583  FILE NTIS
          75  FILE OCEAN
     5144  FILE PASCAL
          2  FILE PHAR
          3  FILE PHARMAML
         18  FILE PHIN
       306  FILE PROMT
          6  FILE PROUSDDR
          9  FILE RDISCLOSURE
     8243  FILE SCISEARCH
     2045  FILE TOXCENTER
     4450  FILE USPATFULL
        361  FILE USPAT2
          8  FILE VETB
         14  FILE VETU
         99  FILE WATER
       718  FILE WPIDS
          3  FILE WPIFV
       718  FILE WPINDEX
L1        QUE PRECONDITIONING
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FILE 'SCISEARCH, BIOSIS, CAPLUS, PASCAL, MEDLINE, USPATFULL, EMBASE,
ESBIOBASE, TOXCENTER' ENTERED AT 14:25:03 ON 09 MAY 2005
L2        1261 S PRECONDITIONING (P) MITOCHONDRIA
L3        508 DUP REM L2 (753 DUPLICATES REMOVED)
L4        15 S L3 AND (ISOCITRATE(W) DEHYDROGENASE(W) NAD(W) ALPHA(W) SUBUNIT O
L5        72 S VANEYK,J?/AU
L6        76 S ARRELL,D?/AU
L7        3875 S ELLIOTT,S?/AU
L8        0 S L5 AND L6 AND L7
L9        6 S EYK,J?/AU
L10       0 S L9 AND L7 AND L8
L11       0 S VANEYK,JENNIFER/AU
L12       2 S VANEYK,JENNIFER?/AU
L13       2 DUP REM L12 (0 DUPLICATES REMOVED)
L14       5 S L6 AND L7
L15       5 DUP REM L14 (0 DUPLICATES REMOVED)

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FILE 'STNGUIDE' ENTERED AT 14:38:33 ON 09 MAY 2005

=>

---Logging off of STN---

=>

Executing the logoff script...

=> LOG Y

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

0.54

134.62

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE

TOTAL

ENTRY

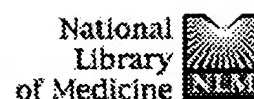
SESSION

CA SUBSCRIBER PRICE

0.00

-1.46

STN INTERNATIONAL LOGOFF AT 14:43:40 ON 09 MAY 2005



All Databases

PubMed

Nucleotide

Protein

Genome

Structure

OMIM

PMC

Journals

Search

PubMed

for

#1 AND (preconditioning)

Preview

Go

Clear

Limits

Preview/Index

History

Clipboard

Details

About Entrez

Text Version

Entrez PubMed

Overview

Help | FAQ

Tutorial

New/Noteworthy

E-Utilities

PubMed Services

Journals Database

MeSH Database

Single Citation Matcher

Batch Citation Matcher

Clinical Queries

Special Queries

LinkOut

My NCBI (Cubby)

Related Resources

Order Documents

NLM Catalog

NLM Gateway

TOXNET

Consumer Health

Clinical Alerts

ClinicalTrials.gov

PubMed Central

- Search History will be lost after eight hours of inactivity.
- To combine searches use # before search number, e.g., #2 AND #6.
- Search numbers may not be continuous; all searches are represented.
- Click on query # to add to strategy

Search	Most Recent Queries	Time	Result
#27	Search #1 AND (preconditioning)	10:41:42	0
#21	Search (isocitrate dehydrogenase NAD alpha subunit OR succinyl CoA ligase OR NADH ubiquinone oxidoreductase OR ATP synthase) AND (preconditioning)	10:34:51	51
#18	Search #7 OR DJ-1 AND (preconditioning)	10:08:34	88
#17	Search #7 OR DJ-1	09:53:30	10586
#11	Search DJ-1	09:51:38	107
#8	Search RNA binding protein regulatory subunit DJ-1	09:36:22	2
#7	Search isocitrate dehydrogenase NAD alpha subunit OR succinyl CoA ligase OR NADH ubiquinone oxidoreductase OR ATP synthase OR prohibitin OR ADP ribosyl hydrolase OR HSP27	09:35:52	10484
#6	Search isocitrate dehydrogenase NAD alpha subunit OR succinyl CoA ligase OR NADH ubiquinone oxidoreductase	09:30:35	3138
#5	Search level of isocitrate dehydrogenase NAD OR succinyl CoA ligase OR NADH ubiquinone oxidoreductase	09:29:15	3281
#4	Search level of isocitrate dehydrogenase NAD alpha subunit	09:27:11	1
#2	Search level of isocitrate dehydrogenase NAD AND alpha subunit	09:24:45	1
#1	Search level of isocitrate dehydrogenase NAD	09:24:22	159

Clear History

Write to the Help Desk

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